### CERTIFICATION

SDG No:

MC45945

Humacao, PR

Laboratory:

Accutest, Massachusetts

Site:

BMS, Building 5 Area, PR

Matrix:

Soil/Groundwater

SUMMARY:

Soil and groundwater samples (Table 1) were collected on the BMSMC facility – Building 5 Area. The BMSMC facility is located in Humacao, PR. Samples were taken May 16, 2016 and were analyzed in Accutest Laboratory of Marlborough, Massachusetts that reported the data under SDG No.: MC45945. Results were validated using the following quality control criteria of the methods employed (MADEP VPH and MAPED EPH, Massachusets Department of Environmental Protection, 2004) and the latest validation guidelines (July, 2015) of the EPA Hazardous Waste Support Section. The analyses performed are shown in Table 1. Individual data review worksheets are enclosed for each target analyte group. The data sample organic data samples summary form shows for analytes results that were qualified.

In summary the results are valid and can be used for decision taking purposes.

Table 1. Samples analyzed and analysis performed

SAMPLE ID	SAMPLE	MATRIX	ANALYSIS PERFORMED
	DESCRIPTION		
MC45945-1	RA-9 GWD	Groundwater	Volatiles TPHC Ranges;
MC45945-1A	RA-9 GWD	Groundwater	Extractable TPHC Ranges
MC45945-2	RA3 (3-4)	Soil	Volatiles TPHC Ranges;
			Extractable TPHC Ranges

Mendez IC # 180

Reviewer Name:

Rafael Infante

Chemist License 1888

Signature:

Date:

June 14, 2016

584967

### **Report of Analysis**

By

AF

Prep Date

n/a

Page 1 of 1

Client Sample ID: RA-9 GWD Lab Sample ID:

MC45945-1

Date Sampled:

n/a

Matrix: Method:

AQ - Ground Water MADEP VPH REV 1.1

DF

1

05/16/16 Date Received: 05/17/16 Percent Solids: n/a

Project:

BMSMC, Building 5 Area, Puerto Rico

Analyzed

05/18/16

**Prep Batch Analytical Batch** 

GBD3638

Run #1 Run #2

Purge Volume

BD73611.D

Run #1 Run #2

5.0 ml

File ID

### **Volatile TPHC Ranges**

CAS No.	Compound	Result	RL	MDL	Units	Q
	C5- C8 Aliphatics (Unadj.)	ND	50	40	ug/l	
	C9- C12 Aliphatics (Unadj.)	ND	50	40	ug/l	
	C9- C10 Aromatics (Unadj.)	ND	50	40	ug/l	
	C5- C8 Aliphatics	ND	50	40	ug/l	
	C9- C12 Aliphatics	ND	50	40	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
	2,3,4-Trifluorotoluene	84%		70-1	30%	
	2,3,4-Trifluorotoluene	100%		70-1	30%	



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

### Report of Analysis

Page 1 of 1

Client Sample ID:

RA-9 GWD

Lab Sample ID:

MC45945-1A

AQ - Ground Water

Matrix: Method: Project:

DF

1

MADEP EPH REV 1.1 SW846 3510C BMSMC, Building 5 Area, Puerto Rico

Date Sampled: 05/16/16 Date Received: 05/17/16

Percent Solids: n/a

Run #1 Run #2 File ID DE14202.D Analyzed 05/21/16

By TA

**Prep Date** 05/18/16

**Prep Batch** OP47537

Analytical Batch

**GDE794** 

**Initial Volume** 

880 ml

**Final Volume** 

Run#1 Run #2

2.0 ml

### **Extractable TPHC Ranges**

CAS No.	Compound	Result	RL	MDL	Units	Q
	C11-C22 Aromatics (Unadj.)	ND	110	80	ug/l	
	C9-C18 Aliphatics	ND	110	80	ug/l	
	C19-C36 Aliphatics	ND	110	80	ug/l	
	C11-C22 Aromatics	ND	110	80	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
84-15-1	o-Terphenyl	80%		40-1	40%	
321-60-8	2-Fluorobiphenyl	67%		40-1	40%	
3386-33-2	1-Chlorooctadecane	47%		40-1	40%	
580-13-2	2-Bromonaphthalene	72%		40-1	40%	

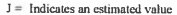


ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range



B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Méndez IC # 1488

### Report of Analysis

Page 1 of 1

Client Sample ID: RA3(3-4)

File ID

16.5 g

AB94125.D

Initial Weight

Lab Sample ID:

MC45945-2

Matrix: Method: SO - Soil

MADEP VPH REV 1.1

DF

1

n/a

Date Sampled: 05/16/16 Date Received: 05/17/16

Percent Solids: 80.7

n/a

Project:

BMSMC, Building 5 Area, Puerto Rico

Analyzed

05/18/16

**Prep Date** 

Prep Batch **Analytical Batch** GAB5173

Run #1 Run #2

Final Volume

16.0 ml

**Methanol Aliquot** 

Run# 2

100 ul

By

DF

Run#1 Run #2

**Volatile TPHC Ranges** 

CAS No.	Compound	Result	RL	MDL	Units	Q
	C5- C8 Aliphatics (Unadj.)	5820	7200	3600	ug/kg	J
	C9- C12 Aliphatics (Unadj.)	145000	7200	3600	ug/kg	
	C9- C10 Aromatics (Unadj.)	79100	7200	3600	ug/kg	
	C5- C8 Aliphatics	5820	7200	3600	ug/kg	J
	C9- C12 Aliphatics	65200	7200	3600	ug/kg	
	-					
	O) O12 / III piaules	03200	7200	3000	og/kg	

CAS No. **Surrogate Recoveries** Run#1

70-130%

Limits

2,3,4-Trifluorotoluene 2,3,4-Trifluorotoluene

81% 83%



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

### **Report of Analysis**

By

TA

Page 1 of 1

Client Sample ID: Lab Sample ID:

RA3(3-4) MC45945-2

SO - Soil

Prep Date

05/24/16

Date Sampled: 05/16/16 05/17/16

Matrix: Method:

MADEP EPH REV 1.1 SW846 3546

Date Received: Percent Solids: 80.7

Project:

BMSMC, Building 5 Area, Puerto Rico

Analyzed

06/02/16

Prep Batch **Analytical Batch** OP47618 GDE803

Run #1

Run #2

**Initial Weight** Final Volume

DF

1

Run #1 Run #2

2.0 ml

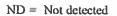
### **Extractable TPHC Ranges**

File ID

11.1 g

DE14424.D

CAS No.	Compound	Result	RL	MDL	Units	Q
	C11-C22 Aromatics (Unadj.)	24200	22000	18000	ug/kg	
	C9-C18 Aliphatics	131000	11000	8900	ug/kg	
	C19-C36 Aliphatics	11100	11000	8900	ug/kg	
	C11-C22 Aromatics	23600	22000	18000	ug/kg	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
84-1 <i>5</i> -1	o-Terphenyl	89%		40-1	40%	
321-60-8	2-Fluorobiphenyl	82%		40-1	40%	
580-13-2	2-Bromonaphthalene	75%		40-1	40%	
3386-33-2	1-Chlorooctadecane	93%		40-1	40%	NO



MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

fael Infant Méndez IC # 1484

N = Indicates presumptive evidence of a compound

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MC45945: Chain of Custody
Page 1 of 3

### **EXECUTIVE NARRATIVE**

SDG No:

MC45945

Laboratory:

**Accutest, Massachusetts** 

Analysis:

MADEP VPH

Number of Samples:

Location:

BMSMC, Building 5 Area

Humacao, PR

**SUMMARY:** 

Two (2) samples were analyzed for Volatiles TPHC Ranges by method MADEP VPH. Samples were validated following the METHOD FOR THE DETERMINATION OF VOLATILE PETROLEUM HYDROCARBONS (VPH) quality control criteria, Massachusetts Department of Environmental Protection, Revision 1.1 (2004). Also the general validation guidelines promulgated by the USEPA Hazardous Wastes Support Section. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

Critical issues:

None

Major:

None

Minor:

None

**Critical findings:** 

None

Major findings:

None

Minor findings:

1. % differences in the rt5.5-7 hydrocarbon range did not meet the method and guidance document performance criteria in the initial calibration verification. No action taken, professional judgment.

2. Continuing and final calibration verification % difference did not meet the hydrocarbon range of rt5.5-7, results were qualified as estimated (UJ) in sample MC45945-2.

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 1888

Signature:

June 14, 2016

Date:

# SAMPLE ORGANIC DATA SAMPLE SUMMARY

Sample ID: MC45945-1

Sample location: BMSMC Building 5 Area

Sampling date: 5/16/2016 Matrix: Groundwater

### METHOD: MADEP VPH

Analyte Name	Result	Units Dilution Factor Lab Flag Validation Reportable	or Lab Flag	Validation	Reportable
Ç5 - C8 Aliphatics (Unadj.)	20	ug/l 1	•	•	Yes
Ç9 - C12 Aliphatics (Unadj.)	20	ug/l 1	•	•	Yes
Ç9 - C10 Aromatics (Unadj.)	20	ug// 1	•	•	Yes
Ç5 - C8 Aliphatics	20	ug/l 1	•	1	Yes
Ç9 - C12 Aliphatics	20	ug/l 1	•	1	Yes

Sample ID: MC45945-2

Sample location: BMSMC Building 5 Area Sampling date: 5/16/2016 Matrix: Soil

## METHOD: MADEP VPH

Analyte Name	Result	Units Dilu	Units Dilution Factor Lab Flag Validation Reportable	Lab Flag	Validation	Reportable
Ç5 - C8 Aliphatics (Unadj.)	5820	ug/kg	П	1	5	Yes
Ç9 - C12 Aliphatics (Unadj.)	145000	ug/kg	₽		⊃	Yes
Ç9 - C10 Aromatics (Unadj.)	79100	ug/kg	1	•	<b>-</b>	Yes
Ç5 - C8 Aliphatics	5820	ug/kg	1	1	D	Yes
Ç9 - C12 Aliphatics	65200	ug/kg	1	1	<b>-</b>	Yes

Type of validation	Full:X Limited:	Project Number:_MC45945
REVIEW OF	VOLATILE PETROLE	UM HYDROCARBON (VPHs) PACKAGE
actions. This document decision and in better according to the data v FOR THE DETER! Massachusetts Depart validation guidelines pr	will assist the reviewer in serving the needs of validation guidance documental omulgated by the USEP of the data	organics were created to delineate required validation using professional judgment to make more informed the data users. The sample results were assessed ments in the following order of precedence METHOD ATILE PETROLEUM HYDROCARBONS (VPH), Protection, Revision 1.1 (2004). Also the general A Hazardous Wastes Support Section. The QC criterian review worksheets are from the primary guidance
The hardcopied (labora has been reviewed and SVOCs included:	atory name) _Accutest_L d the quality control and	aboratories data package received performance data summarized. The data review for
No. of Samples: Field blank No.: Equipment blank No.: Trip blank No.:		Sample matrix:Soil/Groundwater
X Data CompletX Holding TimeN/A GC/MS TuninN/A Internal StandX BlanksX Surrogate ReX Matrix Spike/l	ard Performance	X Laboratory Control SpikesX Field DuplicatesX CalibrationsX Compound IdentificationsX Compound QuantitationX Quantitation Limits
Overall Com (C5_to_C12_Aliphatics	ments:Vola ;_C9_to_C10_Aromatics	tiles_by_GC_by_Method_MADEP_VPH,_REV_1.1
Definition of Qualifiers:		
J- Estimated resu U- Compound not R- Rejected data UJ- Estimated rond Reviewer:	detected	

		Crite	All criteri ria were not met and/d	a were metx or see below
I.	DATA COMPLETNE A. Data Packag			
MISS	SING INFORMATION	DATE LAB. CONTAC	TED DATE	RECEIVED
Ŧ				
В.	Other			Discrepancies:
_0 10	and the second of the second o		5 5	40 55000

All criteria were met	X
Criteria were not met and/or see below	

### HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of extraction, and subsequently from the time of extraction to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE EXTRACTED	DATE ANALYZED	ACTION
Sa	amples analyzed	within method re	commended hold	ing time

### Criteria

### Preservation:

Samples analyzed with ambient purge temperature: Samples must be acidified to a pH of 2.0 or less at the time of collection.

Samples analyzed with heated purge temperature: Samples must be treated to a pH of 11.0 or greater at the time of collection.

Methanol preservation of soil/sediment samples is mandatory. Methanol (purge-and-trap grade) must be added to the sample vial before or immediately after sample collection. In lieu of the in-field preservation of samples with methanol, soil samples may be obtained in specially-designed air tight sampling devices, provided that the samples are extruded and preserved in methanol within 48 hours of collection.

### Holding times:

Aqueous samples using ambient or heated purge - analyze within 14 days. Soil/sediment samples - analysis within 28 days.

Cooler temperature (Criteria: 4 + 2 °C): 2.4°C

Actions: Qualify positive results/nondetects as follows:

If holding times are exceeded, estimate positive results (J) and nondetects (UJ).

If holding times are grossly exceeded, use professional judgment to qualify data. The data reviewer may choose to estimate positive results (J) and rejects nondetects (R).

If samples were not at the proper temperature (> 10°C) or improperly preserved, use professional judgment to qualify the results.

	All crite	eria were metX
	Criteria were not met an	id/or see below
CALIBRATIONS VERIFICATION		
Compliance requirements for satisfactory in that the instrument is capable of producing a		
Date of initial calibration:01/12/16	<u> </u>	02/19/16
Dates of initial calibration verification	:01/12/16	02/19/16
Instrument ID numbers:GC	:AB	GCBD
Matrix/Level:AC	QUEOUS/MEDIUM	

DATE	LAB FILE ID#	ANALYTE	CRITERIA OUT RFs, %RSD, %D, r	SAMPLES AFFECTED
GCAB	<u> </u>	<del></del>		
01/12/16	icv5058-50	rt5.5-7	22.6	None
				,

Note: Initial and initial calibration verification meet method specific requirements except for the cases described in this document. No action taken, professional judgment.

### Criteria- ICAL

- Five point calibration curve.
- The percent relative standard deviation (%RSD) of the calibration factor must be equal to or less than 25% over the working range for the analyte of interest. When this condition is met, linearity through the origin may be assumed, and the average calibration factor is used in lieu of a calibration curve.
- A collective calibration factor must also be established for each hydrocarbon range of
  interest. Calculate the collective CFs for C5-C8 Aliphatic Hydrocarbons and C9-C12
  Aliphatic Hydrocarbons using the FID chromatogram. Calculate the collective CF for
  the C9-C10 Aromatic Hydrocarbons using the PID chromatogram. Tabulate the
  summation of the peak areas of all components in that fraction against the total
  concentration injected. The %RSD of the calibration factor must be equal to or less
  than 25% over the working range for the hydrocarbon range of interest.

### Criteria- CCAL

- At a minimum, the working calibration factor must be verified on each working day, after every 20 samples, and at the end of the analytical sequence by the injection of a mid-level continuing calibration standard to verify instrument performance and linearity.
- If the percent difference (%D) for any analyte varies from the predicted response by more than ±25%, a new five-point calibration must be performed for that analyte. Greater percent differences are permissible for n-nonane. If the %D for n-nonane is greater than 30, note the nonconformance in the case narrative. It should be noted that the %Ds are calculated when CFs are used for the initial calibration and percent drifts are calculated when calibration curves using linear regression are used for the initial calibration.

### Actions:

If %RSD > 25% for target compounds or a correlation coefficient < 0.99, estimate positive results (J) and use professional judgment to qualify nondetects. If % D > 25% (> 30 for nonane), estimate positive results (J) and nondetects (UJ).

### CALIBRATIONS VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:	01/12/16		02/19/16
Dates of continuing calibration	verification:	:05/18/16	05/18/16
Dates of final calibration verific	cation:	05/18/16	05/18/16
Instrument ID numbers:	GCAB_		GCBD
Matrix/Level:	AQUEO	US/MEDIUM	

DATE	LAB FILE ID#	ANALYTE	CRITERIA OUT RFs, %RSD, <u>%D</u> , r	SAMPLES AFFECTED
GCAB	***			
05/18/16	cc5058-50	rt5.5-7	27.7	MC45945-2
05/18/16	cc5058-50	rt5.5-7	28.6	MC45945-2

Note: Continuing and final calibration verification meet method and guidance document specific requirements except for the cases described in this document. Results for hydrocarbon in the range of rt5.5-7 were qualified as estimated (J). Ending calibration verification included in data package.

A separate worksheet should be filled for each initial curve

			Criteria were n	All criteria were met _ ot met and/or see below	
VA. BLANK	ANALYSIS RE	SULTS (Se	ctions 1 & 2)		·-
The assessmen of contaminatio associated with with any blanks determine wheti problem is an is	t of the blank in problems. the samples, s exist, all da her or not the olated occurre amples suspe	analysis res The criteria including tr ita associate ere is an inh	sults is to determing for evaluation of the control of the case of the case of the cata. A secting other data.	ne the existence and mages blanks apply only to disport to disport to disport to the case, of Laboratory Method Blanks in ated to determine if	blanks oblems ated to or if the ok must
List the contam separately.	ination in the	blanks bel	ow. High and low	levels blanks must be	treated
Laboratory biani	ks				
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS	
METHOD BL	ANKS MEET	THE METHO	DD SPECIFIC CRI	TERIA	
Field/Trip/Equip	ment				
				should continually accordively, during sampling, s	
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS	
_NO_TRIP/FIEL	.D/EQUIPMEI	NT_BLANKS	_ASSOCIATED_V	VITH_THIS_DATA_PACI	KAGE.
10 COL P 11			-	1.20	

All criteria were met _	_X
Criteria were not met and/or see below	

### V B. BLANK ANALYSIS RESULTS (Section 3)

### **Blank Actions**

The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. Peaks must not be detected above the Reporting Limit within the retention time window of any analyte of interest. The hydrocarbon ranges must not be detected at a concentration greater than 10% of the most stringent MCP cleanup standard. Specific actions area as follows:

If the concentration is < sample quantitation limit (SQL) and < AL, report the compound as not detected (U) at the SQL.

If the concentration is  $\geq$  SQL but < AL, report the compound as not detected (U) at the reported concentration.

If the concentration is > AL, report the concentration unqualified.

CAMDI E ID

	All criteria were met _	_X
Criteria were no	t met and/or see below	

ACTION

### SURROGATE SPIKE RECOVERIES

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

List the percent recoveries (%Rs) which do not meet the criteria for surrogate recovery. Matrix: solid/aqueous

SUPPOGATE COMPOUND

	2,3,4-Trifluorotoluene		ACTION	SU
RROGATE_ST	ANDARD_RECOVERIES_WITHIN	I_LABORAT	ORY_CONTROL	LIMIT
QC Limits* (Aqu LL_to_L QC Limits* (Sol	JL 70to_130	_to	to	
LL_to_L		to	to	

It is recommended that surrogate standard recoveries be monitored and documented on a continuing basis. At a minimum, when surrogate recovery from a sample, blank, or QC sample is less than 70% or more than 130%, check calculations to locate possible errors, check the fortifying standard solution for degradation, and check changes in instrument performance.

If the cause cannot be determined, reanalyze the sample unless one of the following exceptions applies:

- (1) Obvious interference is present on the chromatogram (e.g., unresolved complex mixture);
- (2) Percent moisture of associated soil/sediment sample is >25% and surrogate recovery is >10%; or
- (3) The surrogate exhibits high recovery and associated target analytes or hydrocarbon ranges are not detected in sample.

If a sample with a surrogate recovery outside of the acceptable range is not reanalyzed based on any of these aforementioned exceptions, this information must be noted on the data report form and discussed in the Executive Report. Analysis of the sample on dilution may diminish matrix-related surrogate recovery problems. This approach can be used as long as the reporting limits to evaluate applicable MCP standards can still be achieved with the dilution. If not, reanalysis without dilution must be performed.

All criteria were met	
Criteria were not met and/or see below	Χ

### VII. A MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples.

At the request of the data user, and in consideration of sample matrices and data quality objectives, matrix spikes and matrix duplicates may be analyzed with every batch of 20 samples or less per matrix.

- Matrix duplicate Matrix duplicates are prepared by analyzing one sample in duplicate. The purpose of the matrix duplicates is to determine the homogeneity of the sample matrix as well as analytical precision. The RPD of detected results in the matrix duplicate samples must not exceed 50 when the results are greater than 5x the reporting limit.
- The desired spiking level is 50% of the highest calibration standard. However, the total concentration in the MS (including the MS and native concentration in the unspiked sample) should not exceed 75% of the highest calibration standard in order for a proper evaluation to be performed. The purpose of the matrix spike is to determine whether the sample matrix contributes bias to the analytical results. The corrected concentrations of each analyte within the matrix spiking solution must be within 70 130% of the true value. Lower recoveries of n-nonane are permissible (if included in the calibration of the C9-C12 aliphatic range), but must be noted in the narrative if <30%.</p>

MS/MSD Recoveries and Precision Criteria

Sample ID:	_MC45945-1	Matrix/Level:_Groundwater/low
Sample ID:	_MC45945-1	Matrix/Level:_Groundwater/low

List the %Rs, RPD of the compounds which do not meet the QC criteria.

The QC reported here applies to the following samples: MC45945-2

Method: MADEP VPH REV 1.1

	MC45958-4a	Spike	MS	MS	Spike	MSD	MSD		Limits
Compound	ug/kg Q	ug/kg	ug/kg	%	ug/kg	ug/kg	%	RPD	Rec/RPD
C5- C8 Aliphatics (Unadj.)	36100	19100	35800	-2* b	19100	36300	1* b	1	70-130/25
C9- C12 Aliphatics (Unadj	.) 762000	21900	684000	-355* b	21900	674000	-401*b	1	70-130/25
C9- C10 Aromatics (Unadj	.) 364000	8210	328000	-439*b	8210	327000	-451*b	0	70-130/25

- (a) Soil to methanol ratio greater than 1.25 to 1.
- (b) Outside control limits due to high level in sample relative to spike amount.

Note: MS/MSD % recoveries outside control limit due to high level in sample relative to amount spike. No action taken.

All criteria were metX	
Criteria were not met and/or see below	

No action is taken on MS/MSD results alone to qualify the entire case. However, used informed professional judgment, the data reviewer may use the MS/MSD results in conjunction with other QC criteria and determine the need for some qualification of the data. In those instances where it can be determined that the results of the MS/MSD affect only the sample spiked, the qualification should be limited to this sample alone. However, it may be determined through the MS/MSD results that the laboratory is having a systematic problem in the analysis of one or more analytes, which affects the associated samples.

### 2. MS/MSD – Unspiked Compounds

List the concentrations of the unspiked compounds and determine the % RSDs of these compounds in the unspiked sample, matrix spike, and matrix spike duplicate.

COMPOUND	CONCENTRAT SAMPLE	TION MS	MSD	%RPD	ACTION
	<u> </u>				
	<del></del>			<del></del>	
3	· 10 20				
				****	

Criteria: None specified, use %RSD ≤ 50 as professional judgment.

Actions:

If the % RSD > 50, qualify the results in the spiked sample as estimate (J). If the % RSD is not calculable (NC) due to nondetect value in the sample, MS, and/or MSD, use professional judgment to qualify sample data.

A separate worksheet should be used for each MS/MSD pair.

All criteria were met Criteria were not met and/or see below _	
MCI E (1.00 (1.00 D) ANNA (1.00 D)	

### VIII. LABORATORY CONTROL SAMPLE (LCS/LCSD) ANALYSIS

This data is generated to determine accuracy of the analytical method for various matrices.

1. LCS Recoveries Criteria

List the %R of compounds which do not meet the criteria

LCS ID	COMPOUND	% R	QC LIMIT	ACTION	
LCS_RE	COVERY_WITHIN_L	ABORATOR'	Y_CONTROL_LIM	rs	
	···			·	

### Criteria:

- \* Refer to QAPP for specific criteria.
- \* The spike recovery must be between 70% and 130%. Lower recoveries of nnonane are permissible (if included in the calibration of the C9-C12 aliphatic range). If the recovery of n-nonane is <30%, note the nonconformance in the executive narrative.

### Actions:

Actions on LCS recovery should be based on both the number of compounds that are outside the %R criteria and the magnitude of the excedance of the criteria.

If the %R of the analyte is > UL, qualify all positive results (j) for the affected analyte in the associated samples and accept nondetects.

If the %R of the analyte is < LL, qualify all positive results (j) and reject (R) nondetects for the affected analyte in the associated samples.

If more than half the compounds in the LCS are not within the required recovery criteria, qualify all positive results as (J) and reject nondetects (R) for all target analyte(s) in the associated samples.

### 2. Frequency Criteria:

Where LCS analyzed at the required frequency and for each matrix (1 per 20 samples per matrix)? Yes or No.

If no, the data may be affected. Use professional judgment to determine the severity of the effect and qualify data accordingly. Discuss any actions below and list the samples affected. Discuss the actions below:

		C	riteria were not met		ia were met ee below _N/A
IX. FIELD/LAE	BORATOR	Y DUPLICATE PR	ECISION		
Sample IDs:	·		Matrix:_		
precision. These a have more varia performance. It is	analyses mability that also expe	neasure both field in laboratory du cted that soil dupli	ken and analyzed as and lab precision; the plicates which mea cate results will have th collecting identical	nerefore asures e a grea	the results may only laboratory ter variance than
COMPOUND	SQL	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
duplicate and	MS/MSD 9	% recoveries RPD	s data package. Blar used to assess preci analytes concentrat	sion. RI	PD within
Criteria:					
RPD + 30% for aq	ueous san	nples, RPD <u>+</u> 50 %	ct-specific informatio for solid samples if r RPD criteria is double	esults a	nre ≥ SQL.
SQL = soil quantita	ation limit				
Actions:					
If both the sample (NC). No action is		uplicate results are	e nondetects (ND), th	ne RPD	is not calculable
Qualify as estimated exceeded the above		ve results (J) an	d nondetects (UJ)	for the	compound that
If one sample resu	ılt is not de	tected and the othe	er is ≥ 5x the SQL qu	alify (J/	UJ).
Note: If SQLs for judgment to determ			are significantly difiate.	ferent,	use professional

If one sample value is not detected and the other is < 5x the SQL, use professional judgment to determine if qualification is appropriate.

All criteria were met _	_X
Criteria were not met and/or see below	

### XI. COMPOUND IDENTIFICATION

The compound identification evaluation is to verify that the laboratory correctly identified target analytes as well as tentatively identified compounds (TICs).

- Verify that the target analytes were within the retention time windows.
  - Retention time windows must be re-established for each Target VPH Analyte each time a new GC column is installed, and must be verified and/or adjusted on a daily basis.
  - o Coelution of the m- and p- xylene isomers is permissible.
  - o All surrogates must be adequately resolved from individual Target Analytes included in the VPH Component Standard.
  - For the purposes of this method, adequate resolution is assumed to be achieved if the height of the valley between two peaks is less than 25% of the average height of the two peaks.
  - The n-pentane (C5) and MTBE peaks must be adequately resolved from any solvent front that may be present on the FID and PID chromatograms, respectively.

Note: Target analytes were within the retention time window.

2. If target analytes and/or TICs were not correctly identified, request that the laboratory resubmit the corrected data.

	Criteria were	e not met and/or see below
XII. QUANTITATION LIN	MITS AND SAMPLE RESULTS	
The sample quantitation eva	aluation is to verify laboratory qu	uantitation results.
1. In the space below,	please show a minimum of one	sample calculation:
MC45945-2	VPH (C7 – C10 Aliphatics)	$RF = 4.015 \times 10^5$
FID		
$[] = (6835351)/(4.015 \times 10^5)$		
[] = 17.02 ppb Ok		
MC45945-2	VPH (C9 – C10 Aromatics)	$RF = 9.58 \times 10^5$
PID		
[] = (1051207456)/(9.58 x 10	5)	
[] = 1,097 ppb Ok		
2. If requested, verify to (MDLs).	nat the results were above the	laboratory method detection limit
3. If dilutions performe the affected samples	d, were the SQLs elevated ac and dilution factor in the table	ccordingly by the laboratory? List below.
SAMPLE ID	DILUTION FACTOR	REASON FOR DILUTION
If dilution was not performed results (J) for the affected co	d and the results were above to impounds. List the affected san	he concentration range, estimate nples/compounds:

All criteria were met \_\_X\_\_\_

### **EXECUTIVE NARRATIVE**

SDG No:

. . . .

MC45945

Laboratory:

**Accutest, Massachusetts** 

Analysis:

MADEP EPH

Number of Samples:

Location:

BMSMC, Building 5 Area

Humacao, PR

SUMMARY:

Two (2) samples were analyzed for Extractable TPHC Ranges by method MADEP EPH. Samples were validated following the METHOD FOR THE DETERMINATION OF EXTRACTABLE PETROLEUM HYDROCARBONS (EPH) quality control criteria, Massachusetts Department of Environmental Protection, Revision 1.1 (2004). Also the general validation guidelines promulgated by the USEPA Hazardous Wastes Support Section. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

**Critical issues:** 

None

Major:

None

Minor:

None

**Critical findings:** 

None

Major findings: Minor findings: None 1. MS/MSD % recovery for C9-C18 Aliphatics above the laboratory control limits in sample

MC45910-3MS/-3MSD. No action taken, MS/MSD results apply only to unspiked sample.

Unspiked sample was from another project.

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 1888

Signature:

Date:

June 14, 2016

### SAMPLE ORGANIC DATA SAMPLE SUMMARY

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Sample ID: MC45945-1A

Sample location: BMSMC Building 5 Area

Sampling date: 5/16/2016
Matrix: Groundwater

METHOD: MADEP EPH

Analyte Name	Result	Units D	ilution Factor	Lab Flag	Validation	Reportable
Ç11 - C22 Aromatics (Unadj.)	110	ug/l	1	-	U	Yes
Ç9 - C18 Aliphatics	110	ug/l	1	-	υ	Yes
Ç19 - C36 Aliphatics	110	ug/l	1	-	U	Yes
C11 - C22 Aromatics	110	ug/l	1	-	υ	Yes

Sample ID: MC45945-2

Sample location: BMSMC Building 5 Area

Sampling date: 5/16/2016

Matrix: Soil

METHOD: MADEP EPH

Analyte Name	Result	Units 1	Dilution Factor	Lab Flag	Validation	Reportable
Ç11 - C22 Aromatics (Unadj.)	24200	ug/kg	1	-	-	Yes
Ç9 - C18 Aliphatics	131000	ug/kg	1		-	Yes
Ç19 - C36 Aliphatics	11100	ug/kg	1		-	Yes
C11 - C22 Aromatics	23600	ug/kg	1	2	2	Yes

Type of validation Full:X Limited:	Project Number:_MC45945
REVIEW OF EXTRACTABLE PETROLE	EUM HYDROCARBON (EPHs) PACKAGE
validation actions. This document will assist the more informed decision and in better serving to were assessed according to the data validation precedence METHOD FOR THE DETERMINATION (VPH), Massachusetts Department (2004). Also the general validation guidelines	le organics were created to delineate required reviewer in using professional judgment to make the needs of the data users. The sample results on guidance documents in the following order of MINATION OF EXTRACTABLE PETROLEUM artment of Environmental Protection, Revision 1.1 promulgated by the USEPA Hazardous Wastes ation actions listed on the data review worksheets to otherwise noted.
The hardcopied (laboratory name) _Accutes received has been reviewed and the quality con review for SVOCs included:	t_Laboratories data package trol and performance data summarized. The data
Lab. Project/SDG No.:MC45945 No. of Samples:2	
X Data CompletenessX Holding TimesN/A GC/MS TuningN/A Internal Standard PerformanceX BlanksX Surrogate RecoveriesX Matrix Spike/Matrix Spike Duplicate	X_ Laboratory Control SpikesX_ Field DuplicatesX_ CalibrationsX_ Compound IdentificationsX_ Compound QuantitationX_ Quantitation Limits
Overall _Extractable_Petroleum_Hydrocarbons_by_GC (C9_to_C36_Aliphatics;_C11_to_C22_(Aromatic	Comments: _by_Method_MADEP_EPH,_REV_1.1
Definition of Qualifiers:	
J- Estimated results U- Compound not detected R- Rejected data UJ- Estimated hondetect  Reviewer:	·

	Criteria were not n	All criteria were metx net and/or see below
I. DATA COMPLETN A. Data Packa		
MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED
	7020 23	
D. Other		
B. Other		Discrepancies:
B. Otner		Discrepancies:
B. Otner		Discrepancies:
B. Other		Discrepancies:
B. Other		Discrepancies:
B. Other		Discrepancies:

All criteria were met	_X
Criteria were not met and/or see below	

### HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of extraction, and subsequently from the time of extraction to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

DATE	DATE	DATE	ACTION
SAMPLED	EXTRACTED	ANALYZED	
- 1 1 1	<u> </u>		
extracted and ar	nalyzed within me	thod recommend	ed holding time
	SAMPLED	SAMPLED EXTRACTED	

### Criteria

### Preservation:

Aqueous samples must be acidified to a pH of 2.0 or less at the time of collection.

Soil samples must be cooled at 4 ± 2 °C immediately after collection.

### Holding times:

Samples must be extracted within 14 days of collection, and analyzed within 40 days of extraction.

Cooler temperature	(Criteria: 4	l <u>+</u> 2 °C):_	2.4°C	
--------------------	--------------	--------------------	-------	--

Actions: Qualify positive results/nondetects as follows:

If holding times are exceeded, estimate positive results (J) and nondetects (UJ). If holding times are grossly exceeded, use professional judgment to qualify data. The data reviewer may choose to estimate positive results (J) and rejects nondetects (R). If samples were not at the proper temperature (> 10°C) or improperly preserved, use professional judgment to qualify the results.

		Crite	All criteria ria were not met and/o	a were metX or see below
CALIBRAT	IONS VERIFIC	ATION		
	at the instrum		nstrument calibration producing and mai	
Dat	e of initial calib	ration:02/04	/16	
Dat	es of initial cali	bration verification:	02/04/13	
Inst	rument ID num	bers:GCD	E	
Mat	rix/Level:	_AQUEOUS/MEDIUN	Λ	
DATE	LAB FILE ID#	ANALYTE	CRITERIA OUT RFs, %RSD, %D, r	SAMPLES AFFECTED
	nitial and conti	auina calibration mo	et method specific requ	iromonto
	initial and Corts	ilung campiation met	st method specific requ	unements

### Criteria- ICAL

- Five point calibration curve.
- The percent relative standard deviation (%RSD) of the calibration factor must be equal to or less than 25% over the working range for the analyte of interest.
   When this condition is met, linearity through the origin may be assumed, and the average calibration factor is used in lieu of a calibration curve.
- A collective calibration factor must also be established for each hydrocarbon range of interest. Calculate the collective CFs for C9-C18 Aliphatic Hydrocarbons, C19-C36 Aliphatic Hydrocarbons, and C11-C22 Aromatic Hydrocarbons using the FID chromatogram. Tabulate the summation of the peak areas of all components in that fraction against the total concentration injected. The %RSD of the calibration factor must be equal to or less than 25% over the working range for the hydrocarbon range of interest.
  - o The area for the surrogates must be subtracted from the area summation of the range in which they elute.
  - The areas associated with naphthalene and 2-methylnaphthalene in the aliphatic range standard must be subtracted from the uncorrected collective C9-C18 Aliphatic Hydrocarbon range area prior to calculating the CF.

### Criteria- CCAL

 At a minimum, the working calibration factor must be verified on each working day, after every 20 samples or every 24 hours (whichever is more frequent), and

- at the end of the analytical sequence by the injection of a mid-level continuing calibration standard to verify instrument performance and linearity.
- If the percent difference (%D) for any analyte varies from the predicted response by more than ±25%, a new five-point calibration must be performed for that analyte. Greater percent differences are permissible for n-nonane. If the %D for n-nonane is greater than 30, note the nonconformance in the case narrative. It should be noted that the %Ds are calculated when CFs are used for the initial calibration and percent drifts are calculated when calibration curves using linear regression are used for the initial calibration.

### Actions:

If %RSD > 25% for target compounds or a correlation coefficient < 0.99, estimate positive results (J) and use professional judgment to qualify nondetects. If % D > 25% (> 30 for nonane), estimate positive results (J) and nondetects (UJ).

### CALIBRATIONS VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:02/04/16						
Dates of co	entinuing calibra	ation verification:_05	/21/16;_06/01/16;_06/	02/16;_06/03/16		
Dates of fin	Dates of final calibration verification:05/21/16;_06/01/16;_06/02/16;_06/03/16					
Instrument	ID numbers:	GCDE				
Matrix/Leve	el:_SOIL/AQUE	OUS/MEDIUM				
DATE	DATE LAB FILE ANALYTE CRITERIA OUT SAMPLES					
	ID#		RFs, %RSD, %D, r	AFFECTED		
Initial and continuing calibration meets method specific requirements. Final calibration verification included in data package.						

A separate worksheet should be filled for each initial curve

				All criteria were met met and/or see below _	
V A DLANIZ	ANALYCIC D				
V A. BLANK	ANALYSIS R	ESULIS (Se	ctions 1 & 2)		
magnitude of complements blanks associated problems with evaluated to decase, or if the	contamination ted with the sany blanks contenting whe problem is armust be run	problems. The samples, inclusives, all data ther or not the isolated occurrence after sample	ne criteria for evaluding trip, equipmand a associated with ere is an inherenturrence not affects suspected of l	etermine the existence uation of blanks apply onent, and laboratory blanks the case must be call variability in the data for the cate of the case must be cast variability in the data for the data. A Laborating other data. A Laborating highly contaminal	only to nks. If refully for the eratory
List the contain separately.	nination in the	e blanks belov	w. High and low l	evels blanks must be ti	reated
Laboratory blar	nks				
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS	
_METHOD BL	ANKS MEET	THE METHO	DD SPECIFIC CR	ITERIA	
Field/Trip/Equip	oment				
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS	
_NO_TRIP/FIE _DATA_PACK/	LD/EQUIPME AGE	NT_BLANKS	S_ANALYZED_AS	SOCIATED_WITH_THI	S

All criteria were met _	_X
Criteria were not met and/or see below	

### V B. BLANK ANALYSIS RESULTS (Section 3)

### Blank Actions

The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. Peaks must not be detected above the Reporting Limit within the retention time window of any analyte of interest. The hydrocarbon ranges must not be detected at a concentration greater than 10% of the most stringent MCP cleanup standard. Specific actions area as follows:

If the concentration is < sample quantitation limit (SQL) and < AL, report the compound as not detected (U) at the SQL.

If the concentration is  $\geq$  SQL but < AL, report the compound as not detected (U) at the reported concentration.

If the concentration is > AL, report the concentration unqualified.

CAMDIEID

All criteria were met _	_X
Criteria were not met and/or see below	

AOTION

### SURROGATE SPIKE RECOVERIES

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

List the percent recoveries (%Rs) which do not meet the criteria for surrogate recovery. Matrix: solid/aqueous

SAMPLE ID	SURROGATE COMPOUND			ACTION		
	S1	S2	S3	<b>S4</b>		
_SURROGATE_ _LIMITS	STANDA	RDS_RECOVE	RIES_WIT	HIN_LABORAT	ORY_CONTROL	
S1 = o-Terphen				fluorobiphenyl		
S3 = 1-Chlorood QC Limits (%)* (			S4 = 2-E	Bromonaphthale	ene 40-140%	
_LL_to_UL_ QC Limits* (Soli	40_to_14		40_to	_14040_to	_140_	
_LL_to_UL_	to	to	to	to		

It is recommended that surrogate standard recoveries be monitored and documented on a continuing basis. At a minimum, when surrogate recovery from a sample, blank, or QC sample is less than 40% or more than 140%, check calculations to locate possible errors, check the fortifying standard solution for degradation, and check changes in instrument performance.

If the cause cannot be determined, reanalyze the sample unless one of the following exceptions applies:

- (1) Obvious interference is present on the chromatogram (e.g., unresolved complex mixture);
- (2) The surrogate exhibits high recovery and associated target analytes or hydrocarbon ranges are not detected in sample.

If a sample with a surrogate recovery outside of the acceptable range is not reanalyzed based on any of these aforementioned exceptions, this information must be noted on the data report form and discussed in the Executive Report. Analysis of the sample on dilution may diminish matrix-related surrogate recovery problems. This approach can be used as long as the reporting limits to evaluate applicable MCP standards can still be achieved with the dilution. If not, reanalysis without dilution must be performed.

All criteria were met	
Criteria were not met and/or see below	_X

### VII. A MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples.

At the request of the data user, and in consideration of sample matrices and data quality objectives, matrix spikes and matrix duplicates may be analyzed with every batch of 20 samples or less per matrix.

- Matrix duplicate Matrix duplicates are prepared by analyzing one sample in duplicate. The purpose of the matrix duplicates is to determine the homogeneity of the sample matrix as well as analytical precision. The RPD of detected results in the matrix duplicate samples must not exceed 50 when the results are greater than 5x the reporting limit.
- The desired spiking level is 50% of the highest calibration standard. However, the total concentration in the MS (including the MS and native concentration in the unspiked sample) should not exceed 75% of the highest calibration standard in order for a proper evaluation to be performed. The purpose of the matrix spike is to determine whether the sample matrix contributes bias to the analytical results. The corrected concentrations of each analyte within the matrix spiking solution must be within 40 140% of the true value. Lower recoveries of n-nonane are permissible but must be noted in the narrative if <30%.</p>

MS/MSD Recov	veries and Precision Crite	eria			
Sample ID:I	MC45910-3		Matrix	/Level:S	oil
List the %Rs, R	PD of the compounds wi	nich do not	meet ti	ne QC criteria.	
MS OR MSD _MC45910-3		% R	RPD	QC LIMITS	ACTION
_MS/MSD	_C9-C18_(Aliphatics)	149/155_	.%	40-140	No_action_

Note: No action taken. MS/MSD results apply only to unspiked sample. Unspiked sample was from another project. No MS/MSD analyzed for aqueous matrix in this data package. No action taken, blank spike/blank spike duplicate used to assess accuracy. % recoveries and RPD within laboratory control limits.

		C	criteria wer	All criteria we not met and/or s	vere metX see below
No action is taken of informed profession conjunction with oth data. In those insta affect only the samp However, it may be a systematic proble associated samples.	al judgment, the er QC criteria and criteria	ne data and deter can be d qualifica ough the l	reviewer imine the determined tion should MS/MSD r	may use the MS need for some qual that the results the limited to this esults that the lab	/MSD results in allification of the of the MS/MSD is sample alone. oratory is having
2. MS/MSD – U	Inspiked Compo	ounds			
List the concentratio compounds in the ur					
COMPOUND	CONCENTRA	ATION MS	MSD	%RPD	ACTION
Criteria: None specif Actions:	ied, use %RSD	≤ 50 as	profession	al judgment.	

If the % RSD > 50, qualify the results in the spiked sample as estimate (J). If the % RSD is not calculable (NC) due to nondetect value in the sample, MS, and/or MSD, use professional judgment to qualify sample data.

A separate worksheet should be used for each MS/MSD pair.

		All criteria were metX Criteria were not met and/or see below				
	VIII.	LABORATORY CONTROL SAMPLE (LCS/LCSD) ANALYSIS				
matric		lata is generated to determine accuracy of the analytical method for various				
	1,:	LCS Recoveries Criteria				
		List the %R of compounds which do not meet the criteria				
LCS II	)	COMPOUND % R QC LIMIT ACTION				
_LCS	S_REC	OVERY_WITHIN_LABORATORY_CONTROL_LIMTS				
	Criteria:  * Refer to QAPP for specific criteria.  * The spike recovery must be between 40% and 140%. Lower recoveries of n-nonane are permissible. If the recovery of n-nonane is <30%, note the nonconformance in the executive narrative. RPD between LCS/LCSD must be < 25%.  Actions:					
		s on LCS recovery should be based on both the number of compounds re outside the %R and RPD criteria and the magnitude of the excedance of teria.				
the as: If the ' for the If more qualify	sociated %R of tagged affected than h	the analyte is > UL, qualify all positive results (j) for the affected analyte in d samples and accept nondetects. The analyte is < LL, qualify all positive results (j) and reject (R) nondetects analyte in the associated samples. The compounds in the LCS are not within the required recovery criteria, sitive results as (J) and reject nondetects (R) for all target analyte(s) in the amples.				
2.	Freque	ency Criteria:				
per ma If no, t the eff	atrix)? <u>Y</u> the data ect and	analyzed at the required frequency and for each matrix (1 per 20 samples <u>'es</u> or No.  The may be affected. Use professional judgment to determine the severity of a qualify data accordingly. Discuss any actions below and list the samples uss the actions below:				

		Crit	All criteria eria were not met and		netN/A below
IX. FIELD/LA	BORATOR	Y DUPLICATE PR	ECISION		
Sample IDs:	<u>.</u>	<u>-</u>	Matrix:		
Field/laboratory duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which measures only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples.					
COMPOUND	SQL	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
	ike duplica	te recoveries RPD	is data package. MS used to assess prec ceptable control limit	ision. R	
Criteria:	should be	raviewed for proje	ect-specific informatio	n	
RPD + 30% for aq	ueous sam	ples, RPD ± 50 %	for solid samples if r RPD criteria is double	esults a	ire ≥ SQL.
SQL = soil quantit	ation limit				
Actions:					
If both the samp calculable (NC). N			s are nondetects (N	D), the	RPD is not
Qualify as estima exceeded the above		e results (J) and	nondetects (UJ) for	the co	mpound that

If one sample result is not detected and the other is  $\geq 5x$  the SQL qualify (J/UJ).

**Note:** If SQLs for the sample and duplicate are significantly different, use professional judgment to determine if qualification is appropriate.

If one sample value is not detected and the other is < 5x the SQL, use professional judgment to determine if qualification is appropriate.

All criteria were met	X
Criteria were not met and/or see below	

### XI. COMPOUND IDENTIFICATION

The compound identification evaluation is to verify that the laboratory correctly identified target analytes as well as tentatively identified compounds (TICs).

- 1. Verify that the target analytes were within the retention time windows.
  - Retention time windows must be re-established for each Target EPH Analyte each time a new GC column is installed, and must be verified and/or adjusted on a daily basis.
  - The n-nonane (n-C9) peak must be adequately resolved from the solvent front of the chromatographic run.
  - o All surrogates must be adequately resolved from the Aliphatic Hydrocarbon and Aromatic Hydrocarbon standards.
  - For the purposes of this method, adequate resolution is assumed to be achieved if the height of the valley between two peaks is less than 25% of the average height of the two peaks.
  - The n-pentane (C5) and MtBE peaks must be adequately resolved from any solvent front that may be present on the FID and PID chromatograms, respectively.
- Aliphatic hydrocarbons range:
  - o Determine the total area count for all peaks eluting 0.1 minutes before the retention time (Rt) for n-C9 and 0.01 minutes before the Rt for n-C19.
  - Determine the total area count for all peaks eluting 0.01 minutes before the Rt for n-C19 and 0.1 minutes after the Rt for n-C36.

Are the aliphatic hydrocarbons range properly determined?

Yes? or No?

Comments:

- 1b. Aromatic hydrocarbons range:
  - Determine the total area count for all peaks eluting 0.1 minutes before the retention time (Rt) for naphthalene and 0.1 minutes after the Rt for benzo(g,h,i)perylene.
  - Determine the peak area count for the sample surrogate (OTP) and fractionation surrogate(s). Subtract these values from the collective area count value.

Are the aliphatic hydrocarbons range properly determined?

Yes? or No?

Comments:

			Criteria were not		vere met; see below _	
2.	If target analytes an laboratory resubmit the			identified,	request that	the
3.	Breakthrough determevaluated for potentially recovery of the fraction and aromatic fraction aphthalene or 2-m the total concentration LCSD, fractionation	al breakthrough actionation sun naphthalene ans of the LCS ethylnaphthaletion for napht	n on a sample sprogate (2-bromornd 2-methylnaph and LCSD. If each in the aliph halene or 2-met	ecific basis ( naphthalene) nthalene in be either the c atic fraction hylnaphtha	by evaluating and on a both the aliphoncentration exceeds 50 lene in the	g the patch hation of the office of the offi
	NOTE:	methylnapht summation	concentration halene in the Long the concestion and the concestion.	CS/LCSD partition de	air includes etected in	the the
	Comments:Concer_concentration_for_n	ntration_in_the aphthalene_ar	_aliphatic_fractio d_2-methylnaph	n_<_5%_of_ thalene	the_total	
	A 10 89					
4.	Fractionation Check containing 14 alkane each constituent. The fractionation efficient optimum hexane volunot allowing signification on the fractionation of the fractionation of the fractional contained in the fractional conta	s and 17 PAH e Fractionation by of each new ame required to ant aromatic he ctionation chec	s at a nominal of Check Solution of lot of silica gel/ defficiently elute ydrocarbon breat k solution, exclu	concentration must be use cartridges, a aliphatic hyd akthrough. F ading n-nona	n of 200 ng/ d to evaluate and establish drocarbons v or each and ane, the Per	ul of the the vhile alyte
	Is a fractionation che	ck standard and	alyzed?		Yes? or No	?
	Comments: Not appli	cable.				

All criteria were met	X
Criteria were not met and/or see below _	

### XII. QUANTITATION LIMITS AND SAMPLE RESULTS

The sample quantitation evaluation is to verify laboratory quantitation results.

In order to demonstrate the absence of aliphatic mass discrimination, the response ratio of C28 to C20 must be at least 0.85. If <0.85, this nonconformance must be noted in the laboratory case narrative.

The chromatograms of Continuing Calibration Standards for aromatics must be reviewed to ensure that there are no obvious signs of mass discrimination.

Is aliphatic mass discrimination observed in the sample?

Yes? or No?

Is aromatic mass discrimination observed in the sample?

Yes? or No?

1. In the space below, please show a minimum of one sample calculation:

MC45945-2

EPH (C11 – C22, Aromatics)

RF = 98200

[] = (10649980)/(98200)

[] = 108.45 ppb Ok

MC45945-

EPH (C19 – C36, Aliphatics)

RF = 66810

[] = (3335834)/(66810)

[] = 49.93 ppb Ok

- 2. If requested, verify that the results were above the laboratory method detection limit (MDLs).
- 3. If dilutions performed, were the SQLs elevated accordingly by the laboratory? List the affected samples and dilution factor in the table below.

SAMPLE ID	DILUTION FACTOR	REASON FOR DILUTION

If dilution was not performed, affected samples/compounds:	(J) for the affect	cted compounds.	List the
			- 1000